

Figure 1 Single lysis solution with Triton X-100. As shown, the tails are preserved and can be stained with dyes for bright field (Wright staining). Chromatin of the halos is also preserved, so it can be well stained with Wright, and the halo edge can be fully viewed and its size accurately established.

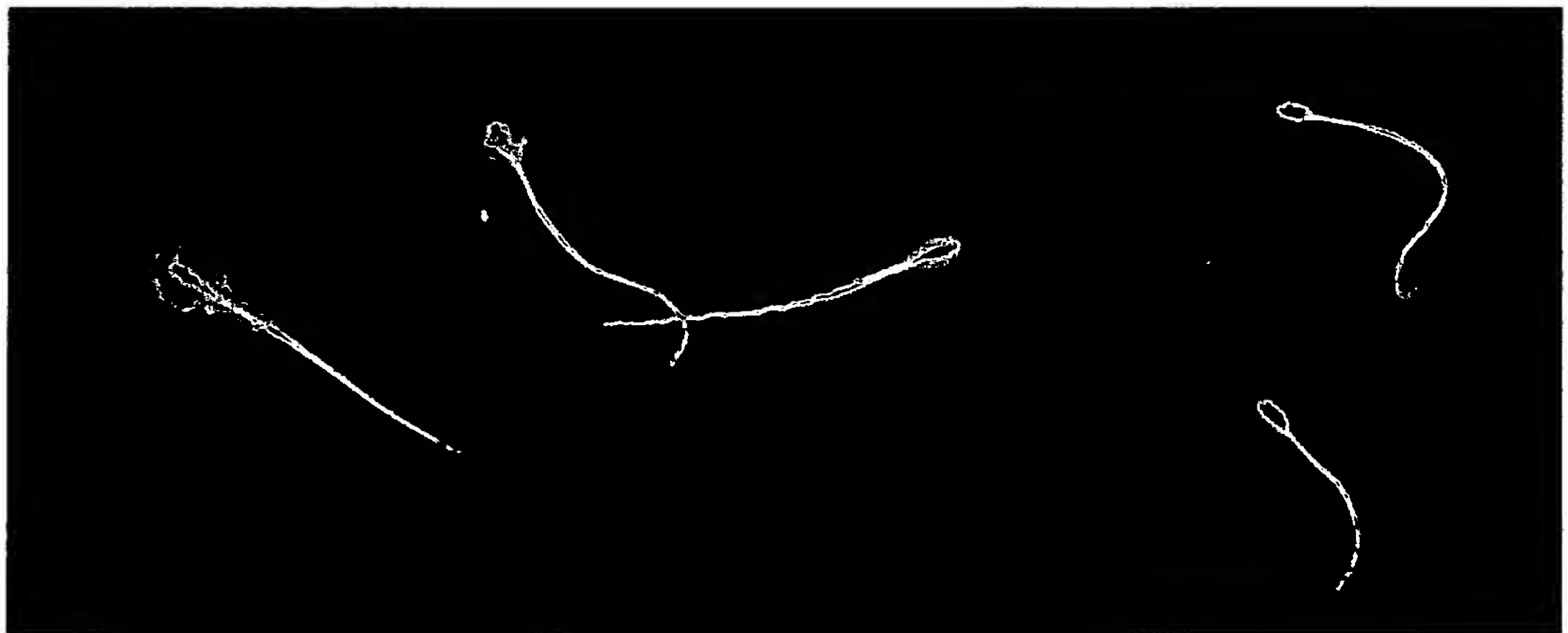


Figure 2 Horse sperm cells. View of the halos of chromatin dispersion and stability of the flagellum after using a single lysis with triton X-100.